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Structural analysis of the N-acetyltransferase Eis1 from Mycobacterium abscessus reveals the molecular determinants of its incapacity to modify aminoglycosides.

**Running title:** Crystal structure of *M. abscessus* Eis1

Kien Lam Ung¹, Laurent Kremer¹,² and Mickaël Blaise*¹

¹Institut de Recherche en Infectiologie de Montpellier (IRIM), Université de Montpellier, CNRS UMR 9004, Montpellier, France.
²INSERM, IRIM, 34293 Montpellier, France.

*Corresponding author:
Tel: (+33)434359447; E-mail: mickael.blaise@irim.cnrs.fr

**Abstract**

Enhanced intracellular survival (Eis) proteins belonging to the superfamily of the GCN5-related N-acetyltransferases play important functions in mycobacterial pathogenesis. In *Mycobacterium tuberculosis*, Eis enhances the intracellular survival of the bacilli in macrophages by modulating the host immune response and is capable to chemically modify and inactivate aminoglycosides. In non-tuberculous mycobacteria (NTM), Eis shares similar functions. However, *Mycobacterium abscessus*, a multi-drug resistant NTM, possesses two functionally distinct Eis homologues, Eis1<sub>Mab</sub> and Eis2<sub>Mab</sub>. While Eis2<sub>Mab</sub> participates in virulence and aminoglycosides resistance, this is not the case for Eis1<sub>Mab</sub>, whose exact biological function remains to be determined. Herein, we show that overexpression of Eis1<sub>Mab</sub> in *M. abscessus* fails to induce resistance to aminoglycosides. To clarify why Eis1<sub>Mab</sub> is unable to modify this class of antibiotics, we solved its crystal structure bound to its cofactor, acetyl-CoA. The structure revealed that Eis1<sub>Mab</sub> has a typical homohexameric Eis-like organization. The structural analysis supported by biochemical approaches demonstrated that while Eis1<sub>Mab</sub> can acetylate small substrates, its active site is too narrow to accommodate aminoglycosides. Comparison with other Eis structures showed that an extended loop between strands 9 and 10 is blocking the access of large substrates to the active site and movement of helices 4 and 5 reduces the volume of the substrate-binding pocket to these compounds in Eis1<sub>Mab</sub>. Overall, this study underscores the molecular determinants explaining functional differences between Eis1<sub>Mab</sub> and Eis2<sub>Mab</sub>, especially those inherent to their capacity to modify aminoglycosides.
**Keywords:** *Mycobacterium abscessus*, *N*-acetyltransferase, aminoglycosides, Eis, GCN5, X-ray structure

**Introduction**

*N*-acetyltransferases are enzymes catalyzing the transfer of acyl groups from acyl-CoA to a wide panel of substrates, ranging from small molecules like spermine to large macromolecules, such as histones. Prokaryotic *N*-acetyltransferases belonging to the GCN5-related *N*-acetyltransferases family (GNAT) are capable to modify a large set of small substrates, including aminoglycoside antibiotics. They can exert pleiotropic effects, participating for instance in the biosynthesis of mycothiol, a mycobacterial antioxidant, or can modify bacterial peptides/proteins as well as host proteins in the case of pathogenic bacteria.

The Eis protein was initially discovered to confer enhanced survival of *Mycobacterium smegmatis* in macrophages, hence its designation as Enhanced Intracellular Survival. Numerous studies have subsequently addressed the function of Eis from *Mycobacterium tuberculosis* (EisMtb) emphasizing its capacity to modulate the host inflammatory response and autophagy as well as its activity to acetylate various proteins, including the host histone H3 and the mycobacterial nucleoid-associated protein HU.

In *M. tuberculosis* clinical strains, mutations in the *eis* promoter resulted in the up-regulation of *eis* expression and resistance to kanamycin. The role of EisMtb in the modification of aminoglycosides has been the subject of deep biochemical and structural investigations, demonstrating that its belongs to the GNAT superfamily of proteins. EisMtb forms a homohexamer and can modify a wide range of aminoglycosides through mono- or poly-acetylations of their primary amine groups.

Consistent with these findings, the Eis2 protein (Eis2Mab) has been shown to also play a key role in *Mycobacterium abscessus*, a non-tuberculous mycobacterium (NTM), recognized as an emerging human pathogen. The *M. abscessus* complex comprises three subspecies, *M. abscessus sensu stricto*, *M. abscessus subsp. bolletii* and *M. abscessus subsp. massiliense* and has been associated with various clinical manifestations, including cutaneous, soft tissues, disseminated infections or lung infections. The pulmonary infections are particularly problematic to treat in the context of cystic fibrosis patients that are highly vulnerable to *M. abscessus* complex. *M. abscessus* is also recognized as one of the most antibiotic-resistant mycobacterial species, leaving clinicians with few therapeutic options. Standard drug regimens against *M. abscessus* lung diseases usually combine three antibiotic classes: an aminoglycoside (amikacin), a macrolide...
(clarithromycin) and a β-lactam (cefoxitin or imipenem)\textsuperscript{14}. However, \textit{M. abscessus} is often refractory to these chemotherapies. The extreme resistance levels of this bacteria to most conventional antibiotics is notably well exemplified by the fact that \textit{M. abscessus} is resistant to all first-line anti-tubercular drugs (isoniazid, rifampicin, pyrazinamide)\textsuperscript{13}.

In addition to drug target modification, resistance mechanisms in \textit{M. abscessus} involve also intrinsic drug resistance, low permeability of the cell wall, induction of drug efflux pumps, absence or non-functional drug-activating enzymes which no longer convert pro-drugs into active metabolites.

Amikacin (AMK), a widely used antibiotic for the treatment of \textit{M. abscessus} lung infection, is inactivated by the \textit{N}-acetyltransferase Eis\textsubscript{2Mab}\textsuperscript{17,18}. Biochemical and structural studies indicated that Eis\textsubscript{2Mab} accommodates a wide range of aminoglycosides in its active site where they are modified by the addition of one or several acetyl groups on their primary amine function, thus impeding their interaction with the ribosome\textsuperscript{18,8}. Solving the high-resolution structure of Eis\textsubscript{2Mab} proved that this protein accommodates amikacin in its active site where the drug undergoes acetylation as supported in enzymatic assays\textsuperscript{18}. Importantly, the deletion of \textit{eis2Mab} increases the susceptibility of \textit{M. abscessus} to aminoglycosides, thus supporting our findings\textsuperscript{17}. Noteworthy, a recent study demonstrated the Eis\textsubscript{2Mab}-dependent enhanced intracellular survival of \textit{M. abscessus} in human macrophages, suggesting that Eis\textsubscript{2Mab} and Eis\textsubscript{Mtb} share a similar biological function\textsuperscript{19}. In contrast to \textit{M. tuberculosis}, the \textit{M. abscessus} genome encodes a second Eis protein homologous to Eis\textsubscript{2Mab}, designated Eis\textsubscript{1Mab} and encoded by \textit{MAB_4124}. However, inactivation of \textit{eis1Mab} only modestly impacted the capacity of the mutant strain to infect macrophages and failed to modify the susceptibility to aminoglycosides as compared to an \textit{eis2Mab} mutant\textsuperscript{19}. This suggests that Eis\textsubscript{1Mab} and Eis\textsubscript{2Mab} share very limited overlapping functions\textsuperscript{17,19}. The reasons for these differential biological functions between Eis\textsubscript{1Mab} and Eis\textsubscript{2Mab} remain obscure when considering the primary sequence identity (31\%) between the two proteins. To fill this gap, and in order to provide the mechanistic basis explaining these differential aminoglycoside-modifying properties, a thorough structural and biochemical study of Eis\textsubscript{1Mab} was undertaken.

\textbf{Materials and Methods}

\textbf{1- Inactivation of \textit{eis1} in \textit{M. abscessus}.}

Deletion of \textit{MAB_4124} (\textit{eis1Mab}) and \textit{MAB_4532c} (\textit{eis2Mab}) in the reference strain \textit{M. abscessus} CIP104536 was carried out using the double homologous recombination unmarked
strategy, reported previously. The DNA sequences located upstream and downstream the eis1Mab gene were PCR-amplified and subsequently cloned into the pUX1-katG using the following primers and containing specific restriction sites (capital letters): fw_up_eis1_NheI 5'-gatcCGCTAGcccgaacccgtcgatacaggagttctcgtc-3'; rv_up_eis1_MfeI 5'-ggatcCAATTGcggtcacggcttcttgccctgtgttgtt-3'; fw_down_eis1_MfeI 5'-gatccCAATTGacttctaggagtctgccgtgtcatccaggaac-3'; rv_down_eis1_HpaI 5'-ggatcGTTAACcaaggctagatgctcgcgcagcgtcgacccg-3'.

The DNA sequences located upstream and downstream the eis2 gene were PCR-amplified and subsequently cloned into the pUX1-katG plasmid using the following primers and containing specific restriction sites (capital letters): fw_up_eis2_HpaI: 5'-catccGTTAACctggagctgctccgatgagccgaaatatgc-3'; rv_up_eis2_MfeI: 5'-gatccCAATTGgtgcgtagggtcagctcagacacatacccatg-3'; fw_down_eis2_NheI: 5'-gatccGCTAGCgggatcgatcagcattactgtgtacagtcc-3'; rv_down_eis2_MfeI: 5'-gatccCAATTGgtttggctggcccgtcgcacccagtgcgcc-3'.

The resulting plasmids were introduced by electroporation in the smooth (S) variant of M. abscessus and the red fluorescent, kanamycin (KAN)-resistant colonies which underwent the first homologous recombination event were selected on Middlebrook 7H10 agar supplemented with 10% Oleic acid-Albumin-Dextrose-Catalase (OADC) in the presence of 200 µg.mL-1 KAN. Cultures of selected colonies were then subjected to a counter-selection on 7H10 agar plates supplemented with OADC containing 50 µg.mL-1 isoniazid (INH) to promote a second homologous recombination and removal of the KAN cassette. Non-fluorescent, INH-resistant colonies were picked up and further genotyped for eis1Mab deletion following PCR using two pairs of primers: fw_up_eis1_NheI with SQ2_b_down: 5'-cttactagaggagccatgggttctc-3' and SQ2_a_down: 5'-cgccaactcctgtgtgggtttggatc-3' with SQ2_b_down. Identification of eis2 mutants was performed by PCR using this pair of primers: SQ1_a_up: 5'-gctgggatgacggcggatgggacgg-3' and SQ1_b_up: 5'-gttggagcgaagatgcgtcag-3'. The positive clones were further confirmed by DNA sequencing of the genomic region surrounding the site of gene deletion.

For functional complementation of the eis1Mab deletion mutant (Δeis1Mab), the genomic sequence of eis1Mab was fused at its 5' end in frame with the sequence encoding the Streptag-II (italic letters), using the following primers (restriction sites in capital letters): fw_Eis1_261_NdeI 5'-ctcaaCATATGatgTGGAGCCACCCGCAAGTTGAGAAAGatgccccctacgcaagcagccatgccc-3' and rv_Eis1_261_HindIII 5'-caatgAAGCTTCgagaattgggtctcagtaaggtgcgcacagcacg-3' and subsequently cloned into pMV261-Zeo in which the kanamycin cassette has been replaced by a zeocine resistance cassette, between NdeI and HindIII. We used similar strategy to complement the Δeis2Mab mutant by cloning the eis2 gene into pMV261-Zeo using the following primers (restriction sites in capital
letters): fw_Eis2_261_NdeI 5’-
ctcaaCATATGatgTGGAGCCACCCGCAGTTCGAGAAGatgtctgagctgaccctacgcaccattg
-3’ and rv_Eis2_261_HindIII 5’-
caatgAAGCTTtcagaagtcgtcgggcgcactgggtgc-
-3’.
The parental, Δeis1 and Δeis2 M. abscessus strains were transformed with either pMV261-Zeo, pMV261-Zeo-eis1 or pMV261-Zeo-eis2 and selected onto 7H10 agar supplemented with OADC and 50 μg.mL⁻¹ Zeocin.

**2-Amikacin susceptibility assessment.**

The different strains were grown in Sauton’s broth medium supplemented with 10%(v/v) OADC/ 0.025%(v/v) tyloxapol and 50 μg.mL⁻¹ zeocin. The exponential bacterial cultures were pelleted and resuspended in PBS buffer. Five microliters of 10-fold serially diluted M. abscessus cultures were spotted on LB plates in the absence or presence of 4, 8 and 12 μg.mL⁻¹ AMK. Plates were incubated at 37°C for 3 days.

**3-Cloning, protein expression, and purification**

The codon-optimized sequence (GenScript) encoding eis₁₅₅₆₉ was cloned into pET-41 between the KpnI and EcoRI restriction sites in frame with the N-terminal Glutathione S-transferase (GST) tag followed by a Tobacco Etch Virus (TEV) protease cleavage site. An overnight culture of *Escherichia coli* BL21 (DE3) strain resistant to Phage T1 (New England Biolabs, Evry, France) carrying pRARE2 and transformed with pET41-eis₁₅₅₆₉ was grown under agitation at 180 rpm at 37 °C in LB medium supplemented with 50 μg.mL⁻¹ KAN and 30 μg.mL⁻¹ chloramphenicol. This preculture was used to inoculate 6 L of LB broth supplemented with the same antibiotics at 37 °C. The cultures were cooled down in ice for 30 min when OD₆₀₀ reached ~0.8 and protein expression was induced with 1 mM of Isopropyl-β-D-thiogalactoside (IPTG) (Euromedex) for 16 h at 18 °C. Cell pellets were collected by centrifugation at 6, 000 g for 20 min and resuspended in buffer A (50 mM Tris-HCl pH 8, 0.4 M NaCl, 1 mM β-mercaptoethanol, and 1 mM benzamidine). Bacteria were lysed by sonication and cell lysates were clarified by centrifugation at 28, 000 g at 4°C for 1 h. The supernatant was then incubated 30 min with GST-sepharose beads with gentle agitation before being loaded onto a gravity column. Beads were washed with 10 column volumes of buffer B (50 mM Tris-HCl pH 8, 0.4 M NaCl, 1 mM β-mercaptoethanol) and buffer C (50 mM Tris-HCl pH 8, 1 M NaCl, 1 mM β-mercaptoethanol). Proteins were finally eluted with buffer D (50 mM Tris-HCl pH 8, 0.2 M NaCl, 1 mM β-mercaptoethanol and 15 mM glutathione). The eluate was incubated with TEV protease (1 mg of TEV protease per 40 mg of protein) and dialyzed overnight against
buffer E (20 mM Tris-HCl pH 8, 0.2 M NaCl, 1 mM β-mercaptoethanol). A Nickel-Nitrilotriacetic acid sepharose column was used to adsorb the His-tagged TEV protease, the cleaved tag and the uncleaved tagged protein. The tag-free protein that weakly binds to the column was collected by washing the column with buffer E supplemented with 50 mM imidazole. The protein was concentrated using a 10 kDa cut-off centricon (Sartorius) and purified by size-exclusion chromatography on a Superdex 200 Increase 10/300 GL column (GE Healthcare) with buffer F (20 mM Tris-HCl pH 8, 0.2 M NaCl). The protein concentration was determined using a Nanodrop 2000c spectrophotometer (Thermo Fisher Scientific) and the protein purity was estimated by Coomassie blue staining. Eis2\textsubscript{Mab} was expressed and purified, as described previously\textsuperscript{18}.

**4-Enzyme kinetic assays**

Eis1\textsubscript{Mab} and Eis2\textsubscript{Mab} were freshly purified in buffer F. The CoA-SH product of the acetyltransferase reaction triggers the reduction of DTNB (5,5-dithio-bis(nitrobenzoic acid) to form TNB (2-thio-5-nitrobenzoic acid, extinction coefficient: 14150 M\textsuperscript{-1}.cm\textsuperscript{-1}) with a peak of absorption at 412 nm. A master stock of KAN (containing 96% Kanamycin A and 4% KAN B), APR (Apramycin), ZEO (Zeocin), AMK (Amikacin), HYG (Hygromycin B), TYR (Tyramine), HIS (Histamine), acetyl-CoA (ACO) and enzymes were dissolved and diluted in buffer F. The premix A (enzyme, ACO, DTNB) and premix B (DTNB, substrate) were first incubated for 5 min at 25 °C. The reaction was initiated and measured immediately after combining both premixes to a final volume of 100 µL containing 0.25 µM enzyme, 0.5 mM ACO, 2 mM DTNB, and 5 mM of each substrate for assessment of the enzyme specificity. For kinetic constants determination one of the concentration either 5 mM TYR or 0.5 mM ACO was kept constant and the concentration of the other one varied. The reaction was followed at 25 °C and data points were recorded every 5 s for 90 s in a quartz cuvette and using a Nanodrop 2000c spectrophotometer (Thermo Fisher Scientific). All absorbance values were normalized against a negative control where the substrate was absent and standard errors were calculated from three replicates. Nonlinear least-squares regression was used to determine the Michaelis-Menten equation (Prism software, GraphPad).

**5-Assessment of protein expression by Western-Blot**

Single colonies were inoculated in 100 mL of Sauton’s broth supplemented with 10%(v/v) OADC/0.025%(v/v) tyloapol and 50 µg.mL\textsuperscript{-1} zeocin and grown for 3 days at 37 °C, under agitation at 120 rpm. Bacteria were harvested by centrifugation and resuspended in buffer A, described above. After opening bacteria by bead-beating and sonication, samples were centrifuged at 20, 000
g, at 4 °C. The supernatant was collected, mixed with loading-blue buffer and separated on SDS-PAGE. Proteins were then transferred onto a nitrocellulose membrane (GE Healthcare) in transfer buffer (20 mM Tris, 150 mM Glycine, 20% (v/v) ethanol). The membrane was saturated by blocking buffer (PBS 1X, 5% (m/v) skimmed milk, 0.2% (v/v) Tween 20) followed by incubation with primary monoclonal Anti-Strep-tag antibodies produced in mouse (dilution 1/10000) (Sigma-Aldrich). Bands were revealed by polyclonal anti-mouse antibodies conjugated to the horseradish peroxidase (dilution 1/5000) (Sigma-Aldrich). Protein were detected by the luminol-based reaction (SuperSignal™—Thermo Fisher Scientific) and imaged by a Chemidoc (Biorad).

6-Crystallization and x-ray data collection

Crystals were grown in sitting drops in MR Crystallization Plates™ (Hampton Research) at 18 °C by mixing 2 µL of protein solution concentrated to 0.4 mg.mL⁻¹ and containing 2 mM ACO with 2 µL of reservoir solution consisting of 0.8 M ammonium sulfate, 0.1 M Bis-Tris pH 5.5 and 1% (w/v) PEG 3350. Crystals were grown for one month and cryoprotected by a brief soaking step into a solution made of 1.4 M ammonium sulfate, 0.1 M Bis-Tris pH 5.5, 1% (w/v) PEG 3350 and 10% glycerol (v/v) prior being cryo-cooled in liquid nitrogen. X-ray diffraction data were collected at the Swiss-light source on the X06DA-PXIII beamline.

7-Structure solving and refinement

The Eis1Mab structure was solved by molecular replacement using the crystal structure of the Eis2Mab monomer bound to ACO (PDB: 6RFT) as a search request and sharing 31 % primary sequence identity with Eis1Mab. Molrep from the CCP4i package 22 was used to performed molecular replacement. Six molecules were found in the asymmetric unit. After a first refinement step leading to R/Rfree of 0.46/0.5, Autobuild from the Phenix software suite 23 was further used to automatically improved the model building which led to R/Rfree: 0.32/0.41. The model was further manually rebuilt and refined using Coot 24 and the Phenix software suite. This led to a final model with refinement statistics shown in Table 1. The coordinates and structures factors have been deposited at the protein data bank under accession number: 6YCA.

Results

1-Overexpression of Eis1Mab does not induce aminoglycoside resistance in M. abscessus
A previous study reported that the deletion of eis1<sub>Mab</sub> but not eis2<sub>Mab</sub>, was associated with an unchanged susceptibility profile to aminoglycosides, leading the authors to conclude that only eis2<sub>Mab</sub> participates in resistance to this class of antibiotics. However, as emphasized by these authors, additional studies are required to confirm or dismiss the implication of Eis1<sub>Mab</sub> in aminoglycoside resistance, particularly by determining the antibiotic susceptibility pattern of a strain overexpressing Eis1<sub>Mab</sub>. To address these two possibilities, eis1<sub>Mab</sub> was cloned into the episomal pMV261-Zeo construct under the control of the strong hsp60 promoter, yielding pMV261-Zeo-eis1, to achieve high levels of protein expression. To easily monitor the expression of Eis1<sub>Mab</sub> by Western blotting, the protein was expressed as a fusion protein harboring a Streptag-II in its N-terminus. Moreover, a Δeis1<sub>Mab</sub> deletion mutant was generated in the smooth variant of M. abscessus using a recently established recombineering procedure that allows the generation of unmarked gene deletion (Figure 1A and Figure 1B). Both the parental and Δeis1<sub>Mab</sub> strains were transformed with either the empty pMV261-Zeo or pMV261-Zeo-eis1. Zeocin was chosen as a selection marker as it does not belong to the aminoglycoside class of antibiotics and thus, the ZEO resistance cassette is very unlikely to affect the aminoglycoside susceptibility profile of the different strains tested. To include a control strain with increased susceptibility to aminoglycosides, we also generated a Δeis2<sub>Mab</sub> deletion mutant known to be susceptible to AMK and complemented strain (Figure S1).

Probing bacterial lysates of three independent clones using anti-Strep tag antibodies testified that high expression levels of Eis1<sub>Mab</sub> and Eis2<sub>Mab</sub> were achieved in both the wild-type, Δeis1<sub>Mab</sub> and Δeis2<sub>Mab</sub> strains carrying pMV261-Zeo-eis1 or pMV261-Zeo-eis2 (Figure 1C and Figure S1). Next, we tested whether overexpression of Eis1<sub>Mab</sub> in both wild-type and Δeis1<sub>Mab</sub> strains was associated with increased resistance to aminoglycosides on agar plates supplemented with increasing concentrations of APR, HYG or AMK. No major growth defect was noticed between the different strains plated on either drug as shown for AMK (Figure 1D). Conversely, the Δeis2<sub>Mab</sub> mutant was hypersensitive to AMK as compare to all the other strains (Figure 1D). Overexpression of Eis2<sub>Mab</sub> in the mutant rescued this effect and growth occurred even in the presence of 12 µg.mL<sup>-1</sup> of AMK, the highest concentration tested (Figure 1D). These results corroborate previous findings suggesting that unlike Eis2<sub>Mab</sub>, Eis1<sub>Mab</sub> is not a primary determinant of resistance to aminoglycosides.

2-Eis1<sub>Mab</sub> is an active N-acetyltransferase but does not modify aminoglycosides
We next investigated the biochemical properties of Eis1Mab. It was previously reported that recombinant Eis1Mab tends to aggregate, rendering the purification procedure challenging. Likewise, we were facing the same issues when Eis1Mab was fused to small tags at its N-terminus, such as His- and S-tags. This tendency to aggregate may suggest a folding issue of the protein and, may explain, at least partly, the lack of activity of Eis1Mab. To circumvent this problem, we attempted to increase protein solubility by overexpressing Eis1Mab fused to the Glutathione-S-transferase (GST) in its N-terminus. This greatly increased the solubility of the protein and to purify it via a three-step purification procedure. Eis1Mab did not aggregate after removal of the GST-tag and, as long it was not too concentrated, a homogeneous and sharp peak could be observed on gel-filtration (Figure 2). The size-exclusion elution profile strongly suggests that Eis1Mab behaves as a multimer in solution. A calibration curve allowed to calculate an apparent molecular weight of 198 kDa, similar to the one reported for its closest homologue Eis2Mab or to those of Eis from M. tuberculosis or M. smegmatis. From this, it can be inferred that Eis1Mab is a hexamer in solution.

The substrate specificity of Eis1Mab was next assessed in vitro by enzymatic assay but no activity was detected towards a range of aminoglycoside antibiotics. However, an N-acetyltransferase activity was detected when using arylalkylamine substrates, such as TYR and HIS (Figure 3A and 3B). These two substrates are smaller compare with aminoglycosides (Figure 3A) and possess primary amine groups. TYR and HIS are known to be acetylated by different N-acetyltransferases and are substrates of Eis from M. tuberculosis and M. smegmatis. Interestingly, Eis1Mab was found to be more active towards TYR and HIS than Eis2Mab (Figure 3B).

This prompted us to determine the kinetic constants of Eis1Mab for TYR, which appears as the preferred substrate of the enzyme (Figure 3B). As could be expected for noncognate substrates the $K_M = 722 \pm 21$ µM and $k_{cat} = 11 \pm 0.5$ s$^{-1}$ for TYR are suggestive of moderate activity. Eis1Mab displays also a slightly lower affinity ($K_M = 144 \pm 19$ µM) for ACO as compared to Eis2Mab ($K_M = 36.2 \pm 2.9$ µM) and to Eis homologous proteins which all display $K_M$ values < 50 µM (Figure 3C). Nonetheless, these in vitro data support the view that Eis1Mab purified under our experimental procedure is an active N-acetyltransferase and, in contrast to Eis2Mab, does not modify aminoglycosides. The smaller substrate preference of Eis1Mab as compared to Eis2Mab makes it a noticeable feature that distinguishes the two enzymes.

3-Overall crystal structure of Eis1Mab

To get insights into the molecular basis driving the Eis1Mab substrate specificity, we aimed at resolving the crystal structure of the protein. Extensive screening for crystallization conditions
using either the apo- or ACO-bound form of Eis1\textsubscript{Mab} generated numerous crystals. However, only crystals of the (ACO)-bound form of Eis1\textsubscript{Mab} allowed the collection of exploitable datasets. We solved the crystal structure by molecular replacement using Eis2\textsubscript{Mab} as a search model and the model was refined to a resolution of 2.9 Å (Table 1). Most residues could be modeled except for the first 23 N-terminus amino acids and part of helix 2 in all monomers.

The overall structure of Eis1\textsubscript{Mab} is a multimer formed by six monomers. This hexameric architecture is similar to the organization of proteins belonging to the GNAT superfamily and also to Eis2\textsubscript{Mab} and to its homologues in \textit{M. tuberculosis} (Eis\textsubscript{Mtb}) and \textit{M. smegmatis} (Eis\textsubscript{Msm}). It consists of the superposition of two homotrimers with a central vestibule (Figure 4A). This oligomeric state fully agrees with the apparent molecular mass determined by size-exclusion chromatography (Figure 2). The structure of the Eis1\textsubscript{Mab} monomer can be divided into three subdomains: the N-terminal GNAT domain (residues 1-135), the central GNAT domain (residues 136-312) and the C-terminal domain (residues 313-415). Each monomer is composed of seventeen β-strands (β1 to β17), ten α-helices (α1 to α10) and seven 3\textsuperscript{10} helices (η1 to η7) (Figure 4B).

4-The Acetyl-CoA binding site of Eis1\textsubscript{Mab}

The best diffracting Eis1\textsubscript{Mab} crystals were obtained in the presence of ACO. The simulated-annealed OMIT electron density map certifies the presence of the cofactor in the Eis1\textsubscript{Mab} structure (Figure 4C). Due to the medium resolution, no water molecule could be modeled but fourteen residues involved in cofactor recognition could be identified (Figure 4D). While the side chains of Leu135 and Ile142 are involved in hydrophobic interactions, the His101 main chain provides a weak H-bond with the oxygen of the acetyl moiety. The carbonyl of Ile102 and Val104 establish also H-bonds. The ACO phosphate groups are recognized by the main chains of Arg110, Gly112, Ile114, and Thr115, as well as by the side chain of Thr115. Arg110 plays a key role in the interaction by making a salt-bridge with the phosphate group, creating a H-bond with the O group and a stacking interaction with the adenine ring of ACO. The carboxyl group of Glu139 binds to the NH2 of the adenine ring whereas Ser138 interacts with the O group of ACO. The Arg145 guanidium group recognizes the O4 of the ribose of ACO (Figure 4D). Additionally, multiple primary sequence alignments indicate that the cofactor binding site is relatively well conserved among the mycobacterial Eis proteins (Figure 5). Therefore, we rule out the possibility that these few amino acid substitutions in the cofactor binding site of Eis1\textsubscript{Mab} would explain the difference of the substrate specificity between Eis1\textsubscript{Mab} and the other Eis proteins.
5- The structural basis for Eis1_Mab substrate specificity

We next searched at the Protein Data Bank using the eFOLD (https://www.ebi.ac.uk/msd-srv/ssm/) and DALI servers for the closest homologues of Eis1_Mab. First of all, we superposed Eis homologues structures from proteins that have been biochemically characterized and have been reported to modify aminoglycosides. The multiple sequence alignments (Figure 5) and superposition of the Eis1_Mab monomer to the other Eis:ACO or Eis:CoA bound structures (Figure 6A) generated the following primary sequence identity and root mean square deviation (r.m.s.d.) values: Eis_Mtb (PDB id: 3RYO, 30% identity, r.m.s.d. 1.8 Å) 30, Eis_Msm (PDB id: 3SXN, 31% identity and r.m.s.d. 1.8 Å) 30, Eis2_Mab (PDB id: 6RFT, 31% seq identity, r.m.s.d. 2.1 Å) 18 and Anabaena variabilis, Eis_Ava, (PDB id: 2OZG, 20% and r.m.s.d. 2.2 Å), unpublished. Additionally, the structure of Eis_Ban from Bacillus anthracis (PDB id: 3N7Z) 29 solved without cofactor displays low sequence identity, 17% but a similar r.s.m.d of 2.1 Å. Our search allowed the comparison of two other Eis homologues that were not biochemically characterized, for instance, the Eis_Efa from Enterococcus faecalis, with no ligand (PDB id: 2I00, 17%, r.m.s.d. 2.10 Å), and Eis_Kfl from Kribbella flavida (PDB id: 4MY0, 21%, r.m.s.d. 2.4 Å). This structural comparison indicate that the closest homologues of Eis1_Mab are Eis proteins from the mycobacterium phylum.

To understand the structural features responsible for Eis1_Mab’s preferences for small substrates, the crystal structure of Eis1_Mab was compared with those of Eis_Mtb and Eis_Msm which were solved in the presence of aminoglycosides. Eis_Mtb was co-crystallized with tobramycin (PDB id: 4JD6) 9 and Eis_Msm with paromomycin (PDB id: 4QB9) 31. Most residues seen interacting with tobramycin in Eis1_Mab are not conserved except for the ultra-conserved carboxy-terminal Phe residue (Figure 5). Furthermore, among the eleven residues contacting paromomycin in Eis_Msm, only three residues are semi-conserved (Thr100, His101, and Glu307) in Eis1_Mab while two others are strictly conserved (Ser138 and Phe415 at the C-terminus) (Figure 5). However, due to the partial conservation in the overall primary sequence between Eis_Mtb, Eis_Msm, and Eis2_Mab (Figure 5), it remains difficult to state that the inability of Eis1_Mab to modify aminoglycosides is inherent to the absence of amino acids conservation in the substrate-binding pocket. We, therefore, hypothesized that the size of Eis1_Mab active site pocket may be reduced as compared with other Eis proteins. Determination of the central cavity volume that includes both the cofactor and substrate binding sites of each Eis protein was carried out with the CASTp server and, using its default settings 32. This analysis showed that Eis1_Mab has a rather small cavity of 651 Å³ in comparison with the other Eis members that were experimentally shown to acetylate aminoglycosides (2005 Å³ for Eis_Mtb, 1678 Å³ for Eis_Msm, 1371 Å³ for Eis_Ava and 1274 Å³ for Eis2_Mab). Only Eis_Ban has a slightly bigger volume than Eis1_Mab (695 Å³) (Figure 6B). Comparison of Eis1_Mab with Eis_Mtb, which possesses the
largest cavity allows proposing that the small cavity in Eis1Mab results of the positioning of helix 4 which pushes further away helix 5. Consequently, this places the bulky side chains of residues Trp205, His210, and Phe213 closer to the Phe415 at the C-terminus, known to be required for substrate recognition (Figure 6C). These structural features reducing the volume of the Eis1Mab substrate cavity are not seen in EisMtb, as the less bulky side chains of the equivalent residues Leu196, Leu200 and Glu202 are not protruding into the aminoglycoside-binding cavity (Figure 6C).

The structural comparison of the various Eis proteins unraveled an atypical feature in the central GNAT domain of Eis1Mab (Figure 5 and Figure 6A). The region composed of fourteen residues and ranging from residues 235 to 248 (235-PQDTAEWFSSSART-248) is forming a loop and a 3_10 helix ($\eta_4$, Figure 4B). This region spanning between strands $\beta$9 and $\beta$10 is extended in Eis1Mab as compared with the other Eis proteins. This loop seems to act as a lid covering the substrate-binding site of the neighbor subunit (Figure 7A). Hence, this region might restrict the accessibility of the substrate-binding pocket to bulky molecules. Furthermore, Phe242, which is not conserved in other Eis homologues (Figure 5) seems of particular interest since its bulky side chain is protruding into the substrate-binding pocket, as can be appreciated when Eis1Mab is superposed onto the tobramycin-bound EisMtb structure (Figure 7A). This may further reduce the volume of the substrate-binding site, thus compromising the access to large substrates, like aminoglycosides. Of note, this loop between strands $\beta$9 and $\beta$10 as well as Phe242 is strictly conserved in all subspecies of the M. abscessus complex (data not shown). Moreover, the corresponding loop was found to be much shorter in all the Eis proteins for which the enzymatic activity towards aminoglycosides was experimentally demonstrated. The loop of EisMtb or EisMas possesses 8 residues (228-VDRTDLKL-235) and 7 residues (226-RGPDGRR-232), respectively (Figure 7B). Additionally, even shorter loops are seen in two other Eis proteins whose activity towards aminoglycosides was demonstrated 29,28 the loops in the EisBan (219-ENYK-222) and EisAva (221-RTRDGS-226) crystal structures are made of 4 and 6 residues, respectively (Figure 7B).

**Discussion**

Mycobacterial N-acetyltransferases and particularly Eis proteins play important roles in macrophage invasion, persistence, modulation of the immune response and antibiotic resistance 33. Eis proteins modify a large set of aminoglycosides and have an impact on antibiotic resistance in numerous mycobacterial species, notably in M. tuberculosis and in NTM, including M. abscessus 18,17 and M. fortuitum 33. Thus, molecules inhibiting Eis proteins given in combination with aminoglycosides would broaden the efficacy of this class of antibiotics in the fight against mycobacterial infections 34. The other biological functions of Eis protein related to the enhanced
intracellular survival or to the modulation of the immune response of the host are also of interest in terms of druggability. Despite being closely related, the contribution of Eis1\textsubscript{Mab} and Eis2\textsubscript{Mab} to the intracellular survival of \textit{M. abscessus} differs. While the deletion of \textit{eis1}\textsubscript{Mab} only slightly impairs the survival of \textit{M. abscessus} in macrophages, the deletion of \textit{eis2}\textsubscript{Mab} profoundly affects the intracellular survival of the mutant strain\textsuperscript{19}. Additionally, regulation of the intracellular expression of the two genes is opposite as \textit{eis1}\textsubscript{Mab} is downregulated while \textit{eis2}\textsubscript{Mab} is upregulated in \textit{M. abscessus}-infected macrophages\textsuperscript{19}. Importantly, our study unambiguously establishes that in contrast to Eis2\textsubscript{Mab}, Eis1\textsubscript{Mab} is not implicated in resistance to aminoglycosides, supporting previous findings\textsuperscript{8,17,35}. This absence of redundant function between the two Eis homologues makes Eis2\textsubscript{Mab} as an interesting drug target to explore further to tackle drug resistance and virulence of \textit{M. abscessus}.

Solving the crystal structure of Eis1\textsubscript{Mab} sheds new light into the differential biological functions of Eis proteins in general and Eis1\textsubscript{Mab} and Eis2\textsubscript{Mab} more particularly. Structural comparison of many Eis structures belonging to the Mycobacterium or to other phyla identified key structural elements that are likely to block the access of Eis1\textsubscript{Mab} binding site to large substrates such as aminoglycosides. In particular, the existence of an extended loop acting as a lid and covering the active site of the neighbor monomer as well as the positioning of helices 4 and 5 restrict the accessibility to the active site. Future work should be dedicated to investigating in more detail the substrate specificity of Eis1\textsubscript{Mab} by creating defined mutants in the extended loop and/or $\alpha$4-$\alpha$5 helices. This appears achievable as modulation of the Eis\textsubscript{Mtb}’s substrate specificity was successfully done by mutating bulky residues that restrict the entrance of the active site\textsuperscript{36}.

This work may also add meaningful information allowing to predict the Eis-like protein substrate specificity. Like \textit{M. abscessus}, other mycobacterial species possess several paralogues of the Eis protein\textsuperscript{33}, and predicting the function of these proteins particularly in the case of pathogenic bacteria may help to anticipate whether these Eis-like proteins may play a role in resistance to aminoglycosides. In that regard, bioinformatics tools such as multiple primary sequence alignments combine with protein structure homology modeling, could be useful to estimate the size of the loop between strands 9 and 10 (or strands 10 and 11 depending of the Eis origin) of the Eis homologues and consequently their substrate specificity.

While Eis1\textsubscript{Mab} cannot modify aminoglycosides, our results indicate that it is an active $N$-acetyltransferase towards the small TYR and HIS arylalkylamine compounds, which are very unlikely the “natural substrates” of the enzyme. In the quest of identifying possible substrate of Eis1\textsubscript{Mab} we carefully analyzed the genomic environment around \textit{eis1}\textsubscript{Mab}, and found that it probably belongs to an operon that is extremely well conserved among all the three \textit{M. abscessus} subspecies and in other NTM, such as \textit{Mycobacterium chelonae}, \textit{Mycobacterium saopaulense} or
*Mycobacterium immunogenum.* This operon also encompasses genes coding for a putative permease (*MAB_4122*), a predicted monoxygenase/acyl-CoA dehydrogenase (*MAB_4123*) located upstream of *eis1*Mab and a nitrilotriacetate monoxygenase (*MAB_4125*) possessing a bacterial luciferase predicted domain, downstream of *eis1*Mab. *MAB_4122* and *MAB_4125* are close homologues of the dibenzothiophene desulfurization enzymes A and C present in numerous bacteria able to degrade dibenzothiophene, an organosulfur compound found in petroleum \(^{37}\). Interestingly, the existence of such degradation pathway has also been described in *Mycobacterium phlei* \(^{38}\), which also possesses one Eis-like protein. Although Eis-like *N*-acetyltransferases have not been found yet functionally linked to such degradation pathway, this conserved genomic organization suggests that *eis1*Mab might be a new partner of this organosulfur chemicals degradation/detoxification pathway. This may be relevant to the fact that NTMs are environmental bacteria, which are likely to encounter organosulfur compounds in their ecological niche. In addition, that dibenzothiophene derivatives are also potent inhibitors of *M. tuberculosis* \(^{39}\) may suggest that the *eis1*Mab operon participates in resistance of *M. abscessus* to these compounds.

**Conflict of interest statement**

The authors declare no conflict of interest.

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**Author contributions**

KLU and MB designed and performed experiments. MB and LK supervised research. MB wrote the first draft with input of all the authors.

**References**


Figure legends

Figure 1: Aminoglycosides susceptibility assessment of Eis1-overexpressing M. abscessus strains.
A Schematic representation of the genomic region around *eis1* in the parental (WT) and Δ*eis1* mutant strains of *M. abscessus*. Sizes of the PCR amplicons used for genotyping of the Δ*eis1* mutant are shown. B PCR profile confirming the proper deletion of *eis1* in the mutant strain. Two PCR products of 1.8 and 2.9 kb were amplified from Δ*eis1* genomic DNA while two bands at 4 and 2.9 kb are expected for the WT strain. C Protein expression pattern of Eis1*Mab* in parental (WT) and knockout strains. Western blotting using anti-Strep-tag primary antibodies attest for the expression of Eis1*Mab* in the two strains transformed with pMV261-Zeo-*eis1* in three individual colonies (C1-C3). Strain transformed with the empty pMV261-Zeo were included as negative controls. The band at 70kDa of the protein ladder possesses also a Strep-tag explaining its immune reactivity with the antibody. D Amikacin susceptibility profile of parental (WT), Δ*eis1* *M. abscessus* strains overexpressing Eis1*Mab* and Δ*eis2* *M. abscessus* strains overexpressing Eis2*Mab*. Serial dilutions of bacteria carrying either pMV261-Zeo, pMV261-Zeo-*eis1* or pMV261-Zeo-*eis2* were spotted onto LB agar supplemented with increasing concentrations of AMK and incubated at 37 °C for 3 days. The figure is representative of two independent experiments.

**Figure 2: Determination of the oligomeric state of Eis1*Mab***

Estimation of the oligomeric state of recombinant Eis1*Mab* by size-exclusion chromatography (left panel). The elution profile of the proteins used for calibration is displayed as a black line and the elution profile of Eis1*Mab* is shown in red. The calibration curve was established using β-amylase (1) (200 kDa), bovine serum albumin (2) (66 kDa), carbonic anhydrase (3) (29 kDa) and cytochrome C (4) (12.4 kDa), eluted with estimated volumes of 11.02, 13.2, 15.8 and 17.4 mL, respectively. The void volume corresponds to the dextran blue elution peak at 8.5 mL. The elution peak of Eis1*Mab* at 10.9 mL matches an apparent molecular weight of 198 kDa. Coomassie Blue-stained SDS polyacrylamide gel electrophoresis of Eis1*Mab* produced after the last step of purification on size-exclusion chromatography (10 µg of protein; right panel).

**Figure 3: Biochemical characterization and enzymatic activity of Eis1*Mab***

A Chemical structures of the different substrates tested in the assay. The structure of zeocin which is a very large molecule is not displayed. B Initial rates of Eis1*Mab* determined in the presence of various substrates, including apramycin, hygromycin, kanamycin, zeocin, tyramine and histamine at a concentration of 5 mM. As a positive control, purified Eis2*Mab* was also tested with TYR and HIS. C Michaelis and Menten curves used to determine the kinetic constants for TYR (left panel) and ACO (right panel).
Figure 4: Overall three-dimensional structure of Eis1\textsubscript{Mab}.

A Overall crystal structure of Eis1\textsubscript{Mab} bound to ACO. The structure of the Eis1\textsubscript{Mab} hexamer is depicted as a cartoon representation where each chain is colored differently. The left panel is a top view of the structure and the right one is rotated by 90°. ACO is pictured in each chain as a sphere representation. B Cross-eye stereo view of the Eis1\textsubscript{Mab} monomer. The three domains are displayed as a cartoon representation where the N-terminal (Nt) GNAT domain is displayed as raspberry color, the central GNAT domain is in blue and the C-terminal (Ct) GNAT domain is in yellow. $\alpha$, $\beta$ and $\eta$ indicate $\alpha$-helices, $\beta$-strands, and 3$_{10}$ helices respectively. C Fo-Fc simulated-annealed OMIT map. The electron density map surrounding the ACO ligand is displayed as a grey mesh and contoured at 2.8 $\sigma$ level. D ACO interactions with the Eis1 residues are shown as white sticks. Hydrogen-bond and the salt-bridge interactions are shown as black dashed lines.

Figure 5: Structural comparison of mycobacterial Eis proteins.

The multiple sequence alignment was performed with ENDscript\textsuperscript{40} and adjusted manually. The secondary structure ($\alpha$, $\alpha$-helix; $\beta$, $\beta$-strand, $\eta$, 3$_{10}$-helix) of Eis1\textsubscript{Mab} extracted from its crystal structure, is indicated above the alignment. Residues involved in ACO binding in Eis1\textsubscript{Mab} are indicated by the green circles, residues involved in aminoglycoside recognition in \textit{M. tuberculosis} (Eis\textsubscript{Mtb}) and \textit{M. smegmatis} (Eis\textsubscript{Msm}) Eis are depicted by the black and cyan square respectively.

Figure 6: Structural comparison of Eis proteins reveals unique features in Eis1\textsubscript{Mab}.

A Superposition of Eis crystal structures. All the structures are displayed as a ribbon representation with the following color codes: Eis1\textsubscript{Mab} in slate, Eis\textsubscript{Mtb} in yellow, Eis2\textsubscript{Mab} in raspberry, Eis\textsubscript{Msm} in magenta, Eis\textsubscript{Ban} in cyan, Eis\textsubscript{Ava} in orange, Eis\textsubscript{Efa} in salmon and Eis\textsubscript{Kfl} in red. Superposition of the carbon $\alpha$ was performed in Coot. The arrow indicates the loop between strands 9 and 10 which appears particularly extended in Eis1\textsubscript{Mab}. B Comparison of the volume (grey surface area) of the active sites of the structures of Eis proteins that have been biochemically characterized, the color code is the same as in A. For structures solved with either ACO or CoA, the cofactor is shown as yellow sticks. C Key structural features restricting the size of the Eis1\textsubscript{Mab} (slate) substrate-binding pocket. Arrows highlight the helices $\alpha\textsubscript{4}$ and $\alpha\textsubscript{5}$ movements seen in Eis1\textsubscript{Mab} as compare with Eis\textsubscript{Mtb} (in yellow). Eis1\textsubscript{Mab} residues side chains that protrude towards the substrate binding-site are shown
as sticks and their equivalent residues in Eis\textsubscript{Mtb} that are not pointing towards the substrate pocket are also indicated.

**Figure 7: Comparison of the extended loop size and conformation**

**A** Close up at the interface of two Eis\textsubscript{1\textsubscript{Mab}} monomers. The extended loop of monomer 2 in slate is covering the active site of monomer 1 (in white) and the Phe242 side chain is pointing towards the substrate-binding site of monomer 1 that is close to the ACO (stick representation) binding pocket. The superposition of Eis\textsubscript{Mtb} structure (PDB id: 4JD6) bound to tobramycin in yellow, to Eis\textsubscript{1\textsubscript{Mab}} highlights the steric hindrance of Phe242, presumably preventing the binding of aminoglycosides. **B** Comparison of the same loop region with the one in other Eis crystal structures from various bacterial origins. Eis\textsubscript{Mtb} is in yellow, Eis\textsubscript{Msm} in magenta, Eis\textsubscript{Ban} in cyan and, Eis\textsubscript{Ava} in orange.
**Figure 1**

(A) Schematic representation of the gene expression pattern in WT and Δeis1 strains. The black bars represent the expression levels of MAB_4123, MAB_4124, and MAB_4125 in WT and Δeis1 strains. (B) PCR analysis showing the expression levels of WT and Δeis1 strains. (C) Western blot analysis showing the expression levels of WT and Δeis1 strains. (D) Colony formation assay showing the effect of AMK on WT and Δeis1 strains.
Figure 2
**FIGURE 3**

A. Chemical structures of APRAMYCIN, HYGROMYCIN B, KANAMYCIN A, AMIKACIN, HISTAMINE, and TYRAMINE.

B. Bar graph showing initial reaction rates with various compounds. **R² = 98.4%** for APR_Eis1, HYG_Eis1, KAN_Eis1, ZEO_Eis, Histamine_Eis1, and **R² = 98.1%** for Tyramine_Eis1 and Tyramine_Eis2.

C. Curves showing initial reaction rates with increasing concentrations of Tyramine and Acetyl-CoA. **R² = 98.4%** for Tyramine, and **R² = 98.1%** for Acetyl-CoA.
FIGURE 4
FIGURE 6

A

B

M. smegmatis

M. tuberculosis

Eis1Mab

M. smegmatis

B. anthracis

Eis2Mab

A. variabilis

C
Supporting information Figure S1: Generation of a Δeis2 M. abscessus mutant and Eis2 overexpressing strain.

A Schematic representation of the genomic region around eis2_Mab in the parental (WT) and Δeis2_Mab mutant strains of M. abscessus. Sizes of the PCR amplicons used for genotyping the Δeis2 mutant are shown. B PCR profile confirming the proper deletion of eis2_Mab in the mutant strain. One PCR product of 1493 bp amplified from Δeis2_Mab mutant genomic DNA is expected while one band at 2657 bp is expected for the WT strain. * Indicates an unspecific amplicon at about 1 kb, systematically obtained with these set of primers using the mutant or WT genomic DNA templates. The dashed lines indicate that two lanes were cropped from the gel. C Protein expression pattern of Eis2_Mab in the parental (WT) and knockout strains. Western blotting using anti-Strep-tag primary antibodies attest for the expression of Eis2_Mab in the two strains transformed with pMV261-Zeo-eis2 in three individual colonies (C1-C3). The strain transformed with the empty pMV261-Zeo was included as a negative control. The band at 70 kDa of the protein ladder possesses also a Strep-tag explaining its immune reactivity with the antibody.